# **EXHIBIT I**

# SUBIM: a program for analysing the Kabat database and determining the variability subgroup of a new immunoglobulin sequence

S.Déret<sup>2</sup>, C.Maissiat, P.Aucouturier and J.Chomilier<sup>1</sup>

#### **Abstract**

Although various programs are available to extract all the information included in protein sequence databases, none is dedicated to immunoglobulins. For this purpose, we designed a program, SUBIM, which is adapted to the Kabat database specialized in immunoglobulin sequences. Besides all the possibilities of any database searching program, SUBIM analyses new sequences of variable regions and determines the variability subgroup they belong to. It also numbers the new sequence according to the system established by Kabat and co-workers for an easier comparison with the other immunoglobulins, thus realizing an automatic alignment with other members of a given type of immunoglobulin chain. This program is largely machine independent and requires very little memory, and should help biochemists concerned with new immunoglobulin sequences.

#### Introduction

The current accumulation of immunoglobulin (Ig) sequences, and the need for comparison studies in correlation with Ig binding specificities or pathogenic properties, require the use of an exhaustive and performing dedicated database. The Kabat database (Kabat et al., 1991) is presently the most complete collection of amino acid sequences of Ig and related proteins of immunological interest. The 5.0 release of Kabat database contains 469 human heavy, kappa and lambda chain variable (V) regions  $(V_H, V_{\kappa} \text{ and } V_{\lambda} \text{ respectively})$  compared to the 132 found in SwissProt release 29.0 (Bairoch and Boeckman, 1991) by retrieving all files containing the keywords immunoglobulin' and human' with the Genetics Computer Group (GCG) method (Devereux et al., 1984). All human Ig sequences from SwissProt are included in the Kabat database, but the  $V_H^{\prime}$ ,  $V_{\kappa}$  and  $V_{\lambda}$  of SwissProt correspond only to 26%, 29% and 33% respectively of those collected by Kabat and coworkers when taking into account sequences of 35 residues or more. In addition, many small pieces of sequences are

included in the Kabat database. These are of interest because the original numbering allows all V regions of a given type  $(\kappa, \lambda \text{ or } H)$  to be directly aligned, so their position can be located within the full sequence. This numbering takes into account the frequent amino acid deletions and insertions which occur at certain positions as a result of genetic divergence of the V gene segments and junctionnal variability during the somatic rearrangements of Ig genes.

To our knowledge, no dedicated software has been developed to use efficiently the Kabat database. In this paper, we present the program SUBIM which allows selection and extraction of V and constant (C) region amino acid sequences for comparison studies. It also allows these sequences to be numbered according to Kabat methodology. Compared to other available programs, SUBIM clusters Ig V regions into sets of related sequences which are called variability subgroups. Since the early studies of Milstein (1967) and Hilschmann (1969), subgroup classification has become (mainly by the monumentous work of Kabat and colleagues) a guiding principle of structural immunology. This taxonomy is based upon the conservation of residues between pairs of sequences and within one subgroup. For example, the mean sequence identities within subgroups  $\kappa II$  and  $\lambda VI$ are close to 80% in both cases, with a root mean square deviation of  $\sim$ 5, while the identity between their consensus is 44%. The identity between the consensus sequences of the four  $\kappa$  subgroups is between 61 and 74%. For subgroups I-IV of human  $\alpha$  chains, this classification was fully confirmed when the human germline sequences on one allele became known (Zachau and Schäble, 1993).

#### Algorithm

Extraction of information concerning one or a set of sequences

A first use of SUBIM is for scanning the database, using keywords that can be selected by any combination based on Boolean algebra.

Allowed keywords are:
protein name (full or partial)
species
author
year of publication

Laboratoire d'Immunologie et Immunopathologie, CNRS URA 1172, F-86021 Poitiers and <sup>1</sup>Laboratoire de Minéralogie-Cristallographie, Universités Paris VI et VII, CNRS URA 009, Case 115, F-75252 Paris Cédex 05, France

<sup>&</sup>lt;sup>2</sup>To whom correspondence should be addressed. Email: criso@zeus.univpoitiers.fr

isotype and variability subgroup domain.

All this information can be retrieved from the database as long as these fields are documented in the files. Another possibility is retrieving information within sequences, such as selecting all Ig sequences with a given residue at a determined position.

Different output formats for the retrieval of a sequence are possible. An example of a request with keywords through SUBIM is presented on Figure 1, with the short

#### 

- 1: Selection
- 2: New sequence
- 3: Listing of the selection
- 4 : End

#### Enter your choice: 1

#### ####### SELECTION MENU #######

- 1: by name
- 2: by species
- 3: by authors
- 4: by year of publication
- 5: by isotype
- 6: by domain
- 7: by residue at precise position
- 8: reset selection
- 9: save the selection
- 10: make the selection
- 11: quit

#### Enter your choice :2

several selections are successively performed until the option 10 is chosen to finish the process of selection

Enter your choice:10

#### Selection Request

(SOURCE = HUMAN) AND

(ISOTYPE = KAPPA LIGHT) AND

(POS = 30 RES = A) AND

#### >>> There are 2 proteins in the selection set

#### 

- 1: short output of selected proteins
- 2: full Kabat output of selected proteins
- 3: save short output
- 4: save full output
- 5 : sequence alignment of selected proteins
- 6 : quit

#### Enter your choice: 1

KEA HUMAN Kappa Light chain Variable Subgroup 3 SHM HUMAN Kappa Light chain Variable Subgroup 3

Fig. 1. Systematic research. Monoclonal Ig light chains of the  $V_{\kappa I}$  subgroup involved in Fanconi's syndrome have been sequenced. Two of them (CHEB and TRE) include one residue that has never been described before in this subgroup: Ala at position 30. Thus, it is worth looking for this residue in every monoclonal light chain which might cause Fanconi's syndrome. This figure shows the succession of menus and the short output option. Ala 30 already exists in two human  $\kappa$  chains of the  $V_{\kappa III}$  subgroup, KEA and SHM.

output option. This short output only gives the protein name, species, isotype and variability subgroup. If this reduced information is insufficient, a longer output is possible, where all the information available in the database is retrieved, including the sequence and all comments. An example of a long output is shown in Figure 2.

Determination of the variability subgroup of a new sequence

Because of the high homology within one subgroup, especially in the N-terminal part, the first 15 residues are sufficient to determine the variability subgroup. The N terminus is a rather accessible data even when small amounts of the purified protein are available, while the complete amino acid sequence may remain undetermined in many instances.

Consensus sequences within each subgroup were previously determined by Kabat et al. (1991). Comparisons

ENTRY HKL325 #Type Protein

TITLE Kappa light chain subgroup III V region (KEA) Human

DATE 15-Oct-1991

PLACEMENT 29.0 78.0 0.0 0.0 0.0

SOURCE #Common-name human

ACCESSION A11851

#### REFERENCE

#Authors Wang A.C., Fudenberg H.H.

#Journal Immunol. Commun. (1975) 4:483-497

#Reference-number R10919

#Comment Checked by author 23-Sep-1977

#### REFERENCE

#Authors Wang A.C., Tung E., Wang I., Fudenberg H.H., Pick

A.I., Froehlichman R.

#Journal Cancer immunol. (1980) 9:81-86

#Reference-number R10890

#Comment Checked by author 18-Mar-1981

#### **GROUP-CODE HKL3**

SUMMARY #Molecular-weight #Length 40 #Checksum 3969 SEQUENCE

5 10 15 20 25 30

#### 1 EIVLTQSPATLSLSPGERATLSCRAGSBVA

#### 31 K S L A W Y Z Z K P

///

Fig. 2. Example of long output format. The long output option of the print menu allows the user to retrieve all information available in the database on both human  $\kappa$  chains of the  $V_{\kappa III}$  subgroup, described in Figure 1. The figure only gives the example of KEA.

with two consensus sequences of each subgroup (4  $\kappa$  and 6  $\lambda$  subgroups) proved sufficient in all studied cases. The first consensus corresponds to the most common residue at each position, and the second consensus corresponds to the second highest occurrence at each position. A frequency value (F) is attributed to each residue of the consensus sequences:

F = occurrence of this residue at this position/ number of sequences in the subgroup

A residue with a frequency of between 0.95 and 1.0 is considered 'invariant'.

The 15 N-terminal residues of a new sequence are compared with the 2N consensus, N being the number of subgroups. If the first amino acid matches that in one consensus, a S score attached to that position is given the

```
query sequence:
   DIVMTQSPESLAVSL
\frac{\text{first consensus sequence kappa I (most frequent):}}{X \ D \ I \ Q \ M \ T \ Q \ S \ P \ S \ S \ L \ S \ A \ S \ V \ G \ D \ V \ R \ T}
0.016 0.748 0.759 0.710 0.721 0.819 0.803 0.819 0.814 0.776 0.579
0.841 \ \ 0.879 \ \ 0.672 \ \ 0.798 \ \ 0.699 \ \ 0.819 \ \ 0.639 \ \ 0.743 \ \ 0.683 \ \ 0.803
\sum F_{\text{first consensus kappa}} = 15.14
F at each position:
0.005 0.022 0.016 0.022 0.087 0.005 0.038 0.005 0.005 0.011 0.289
0.033 \quad 0.022 \quad 0.093 \quad 0.022 \quad 0.120 \quad 0.005 \quad 0.109 \quad 0.027 \quad 0.114 \quad 0.032
S1 = 0.748 + 0.759 + 0 + 0.721 + 0.819 + 0.803 + 0.819 + 0.814 + 0 + 0.579 + 0.841
    +0+0+0.798+0/15.14=0.508
S2 = 0 + 0 + 0 + 0.005 + 0 + 0 + 0 + 0 + 0 + 0 + 0 + 0 + 0.672 + 0 + 0 + 0 / 15.14 = 0.049.
S3 = 0 + 0 + 0 + 0 + 0 + 0 + 0 + 0.776 + 0 + 0 + 0.672 + 0 + 0 + 0.699 + 0 + 0 / 15.14
  = 0.141.
S5 = 0 + 0 + 0 + 0 + 0 + 0.011 + 0 + 0 + 0.093 + 0 + 0.798 + 0 + 0 + 0 + 0.027 + 0 / 15.14
+0+0/15.14=0.126.
Smax (for kappa I) = 50.8\%
```

Fig. 3. Determination of subgroup variability of a new sequence. A new N-terminal sequence of a light chain responsible for non-amyloid lightchain deposition disease was determined. For each consensus of the subgroup  $V_{\kappa I}$ , the frequency F of every 15 N-terminal residue is given, which is the probability of finding such a residue at this position in the subgroup. X indicates that several amino acids were found at this position, with none being found significantly more frequently than others. SUBIM compares this sequence with each consensus sequence of the  $\kappa I$  variability subgroup. The S0 score is determined by summing the 15 values of F when a match is observed between the query and one of both consensus. If no match is observed, no value is added. The query sequence is shifted by 1, hence residue D becomes residue 2, and the S1 score is calculated. Seven scores are calculated by successive shifts. This operation is repeated for each variability subgroup. The sequence is finally assigned to the  $\kappa$  light-chain subgroup IV, which yields the highest score (73.1%).

Smax (for kappa II) = 50.7%

Smax (for kappa III) = 41.8%

Smax (for kappa IV) = 73.1%

frequency of the corresponding residue in this consensus. At the second position an equivalent value is added to this S score, and so on until the 15th residue. The result is divided by the sum of first consensus frequencies, leading to a S0 score.

In a second step, the query sequence is shifted by 1, and hence starts at position 2. A new score is then determined similarly. Six successive shifts are performed leading to scores S0-S6. The limit of six residue shifts is based on the fact that an Ig sequence may be truncated by at most six amino acids at the N-terminal part, when aligned with the family. This value is acertained from the fact that the starting of numbering, within any subgroup of the Kabat database, is included in the range -6, +6, whatever sequence is considered. This was verified for 98.7% of the sequences.

Finally seven scores are determined for each subgroup and the highest value allows a subgroup to be attributed to the new sequence.

#### Numbering new sequences of V regions

In the Kabat database, the V region sequences are aligned to permit various comparisons of the  $V_{\kappa}$  and  $V_{\lambda}$  from different species (human, mouse, etc.) and different variability subgroups. This was performed by sequential numbering with gaps inserted for alignment. Most of these gaps are located in the complementarity determining regions (CDRs). Tramontano and Lesk (1992) demonstrated that the Kabat numbering is required for comparing Ig chains.

Strictly conserved residues within one isotype  $(\kappa, \lambda)$  or heavy chain) may be used as anchor points since they are particularly important for the structure and functions. Such highly conserved residues are encountered at several positions spread all over the sequences and in a cluster at the beginning of the third framework (FR3) segment.

First, SUBIM searches for known invariant residues (e.g. Cys23, Trp35, Gly57, Cys88 in a  $\kappa$  chain) in the new sequence and places them at the corresponding positions. It then calculates the number of amino acids between these conserved residues. According to this difference value, gaps are filled if necessary, according to the flexible Kabat notation. For instance, if the residues numbered 27 and 28 in Kabat numbering are also found in the new sequence but separated by two residues, a gap is introduced with numbers 27A and 27B. This format procedure allows alignment of new determined amino acid sequences with sequences already available in the Kabat database. It was not developed for C domains because their sequences are highly conserved.

Figure 4 shows an example of new Ig, whose sequences are not yet in the Kabat database; the subgroup of each

#### 

- 1: short output of selected proteins
- 2: full Kabat output of selected proteins
- 3: save short output
  4: save full output
- 5: sequence alignment of selected proteins
- 6 : quit

Enter your choice: 5

#### **ALIGNED SEQUENCES:**

REI										10										20	)
	D	I	Q	M	T	Q	S	P	S		L	S	Α	S	٧	G	D	R	V	T	
CHEB	D	I	Q	M	T	Q Q	S	P P	S	S S	L	S	Α		V	G	D	R	V	T	
TRE	D	i	Q	Μ	T	Ò	S S S	P	S	S	L	S	Α	S	V			R	V		
			Ì		•	•															
							27	A	В	C	D	E	F	28		30					
REI	I	T	С	Q	Α	S	Q	-	•	•	-	-	-	D	I	I	K	Y	L	N	
CHEB	F	T	С	R	Α	S	Q	-	-	-	-	•	-	T	I	Α	T	F	L	N	
TRE	1	T	C	<u>R</u>	Α	S	Q	-	-	-	-	•	-	S	I	<u>A</u>	G	Y	L	<u>N</u>	
											Cl	DRI									
						40										50					
REI	W	Y	Q	Q	T	P	G	K	Α	P	K	L	L	I	Y	E	Α	S	N	L	
CHEB		Ŷ	ŏ		ĸ	P	G	K	A	P	K	L	Ĺ	i	Ŷ	Ğ	A	Š	S	L	
TRE		Ŷ	õ	õ	R	P	Ğ	ĸ	A	P	E	Ľ	Ĺ	Î	Ŷ	Ť	A	Š	Ť	Ĺ	
		-	•	•	••	•	_	••	••	•	-	_	_	•	-	_			)R2		
																		-			
						60										70					
REI	Q	Α	G	v	P	S	R	F	s	G	s	G	s	G	Т	D	Y	T	F	T	
CHEB	Q	S	G	V	P	S S	R	F	S	G	S	G	S	G	T	D D	Y F	T	L	T	
		S	G	V		S	R R R	F F	S S S	G G G	S S		S			D	Y				
CHEB	Q	S	G	V	P	S S	R	F	S	G	S	G	S	G	T	D D D	Y F	T	L	T	
CHEB TRE	Q R	\$ <u>\$</u>	G G	V V	P P	S S S	R R	F	S S	G G	S S	G G	S S	G G	T A	D D	Y F F	T	L L	T T	
CHEB TRE	Q R I	\$ <u>\$</u> \$	G G S	V V L	P P	S S S 80 P	R R	F F D	S S	G G A	S S	G G Y	S S	G G C	T A Q	D D Q	Y F F	T T	L L S	T T L	
CHEB TRE REI CHEB	Q R I I	\$ \$ \$	G G S	V V L L	P P Q Q	S S S 80 P P	R R E E	F F D	S S I F	G G A A	S S T T	G G Y Y	S S Y Y	G G C C	T A Q Q	D D D · 99 Q Q	Y F F	T T Q Y	L L S S	T T	
CHEB TRE	Q R I	\$ <u>\$</u> \$	G G	V V L	P P	S S S 80 P	R R	F F D	S S	G G A A	S S	G G Y Y	S S	G G C C	T A Q	D D D · 99 Q Q	Y F F	T T	L L S	T T L	
CHEB TRE REI CHEB TRE	Q R I I I	\$ \$ \$	G G S	V V L L	P P Q Q	S S S 80 P P	R E E E F	F F D D D	S S I F S	G A A A	S S T T	G G Y Y Y	S S Y Y Y	G G C C C	T A Q Q Q	D D D · 99 Q Q	Y F F Y S	T T Q Y Y	L L S S	T T L I Y	
CHEB TRE REI CHEB TRE	Q R I I I 95 P	\$ \$ \$ \$ \$	G G S S	V V L L L	P P Q Q	S S 80 P P	R R E E	F F D D D Y	S S I F S	G A A A	S S T T T	G G Y Y Y 100 Q	S S Y Y Y	G C C C	T A Q Q Q	D D . 90 Q Q Q L	Y F F Y S S	T T Q Y Y	L L S S	T T L I Y	R
CHEB TRE  REI CHEB TRE  REI CHEB	Q R I I I 95 P	S S S S	G G S S	V V L L L	P P Q Q	S S 80 P P	R E E E F	F F D D D 96 Y W	S S I F S	G A A A F F	S S T T T G	G G Y Y Y 100 Q	S S Y Y Y G G	G C C C T T	T A Q Q Q K K	DDD . 90 QQQ LV	Y F F Y S S	T T Q Y Y I I	L L S S S	T T L I Y 107 T K	R
CHEB TRE REI CHEB TRE	Q R I I I 95 P	S S S S	G G S S	V V L L L	P P Q Q	S S S S P P P E	R E E E F	F F D D D Y	S S I F S	G A A A	S S T T T	G G Y Y Y 100 Q	S S Y Y Y	G C C C	T A Q Q Q	D D . 90 Q Q Q L	Y F F Y S S	T T Q Y Y	L L S S S	T T L I Y	

Fig. 4. Example of sequence alignment. TRE and CHEB, two  $V_{\kappa I}$  Ig light chains involved in Fanconi's syndrome, are aligned with REI, a  $V_{\kappa}$  dimer whose three-dimensional structure is known. The sequence identity is 77% between TRE and REI, and 82% between CHEB and REI.

sequence has been determined and the numbering performed, allowing alignment with REI, a  $V_{\kappa}$  light chain dimer whose three-dimensional structure is available in the Protein Data Bank (Bernstein *et al.*, 1977). This alignment is a first step in the procedure of building a three-dimensional model by homology modelling.

#### Implementation

SUBIM has been written in the C programming language and has been implemented under the Unix operating system. The program source code is available by anonymous ftp at the node ftp.lmcp.jussieu.fr, under the pub directory. Memory requirements for the microcomputer are modest, but a hard disk with sufficient capacity to hold the Kabat database is necessary.

#### **Discussion**

We have developed a tool for exploiting the currently most complete database for Ig sequences at the protein level. SRS (Etzold *et al.*, 1993), an information indexing and retrieval system designed for libraries such as the EMBL

nucleotide sequence databank (Stoehr and Cameron, 1991), the SwissProt protein (Bairoch and Boeckman, 1991) and the PROSITE library of protein subsequence consensus patterns (Bairoch, 1991) may also perform most SUBIM functions. SRS allows exploitation of all information provided by these libraries (Etzold and Argos, 1993a,b), although some knowledge of the libraries is essential for its use. Elgavish and Schroeder (1993) recently developed a program, SAW, designed for the analysis of Ig V region nucleotide sequences; it is especially interesting for comparing complementary DNA sequences extracted from GenBank or standard text files, assigning functionnal Ig V regions to known germline segments, translation to amino acid sequences and for graphical presentation of data. Our program is dedicated to the exploitation of the Ig amino acid sequences, and is thus complementary to SAW.

SUBIM was designed not only for searches but also for some useful functions, such as V region variability subgroup determination, identification of unusual residues, and numbering according to Kabat et al. (1991). SUBIM allows the user to extract and study known human heavy- and light-chain sequences. For instance, comparisons of primary structures of monoclonal Ig chains may be particulary relevant to the understanding of their pathogenicity (Preud'homme et al., 1994). It may be also interesting to look for precise residues at certain positions, which may be related to an unusual behavior of an Ig chain: for instance, an asparagine at position 70 of a  $\kappa$  chain generally creates a functionnal N-glycosylation site that may cause myeloma complications such as lightchain deposition disease (Cogné et al., 1991) or amyloidosis (Aucouturier et al., 1992). Determining the variability subgroups may be useful because some subgroups are more frequently expressed in some complications of light-chain myeloma (Solomon et al., 1982; Denoroy et al., 1994).

At this time, procedures for determining variability subgroups of new sequences and formatting them, run only for human sequences but extension to other species is currently in progress.

In the last version of the Kabat database (1994), sequences are clustered by families, based on amino acid sequence only. Each family is composed of sequences that differ by 12 amino acids or less. Thus, in addition to other uses, SUBIM may be valuable in preserving the subgroup clustering.

#### Acknowledgements

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